On the Presence of a Proteolytic Enzyme in Casein

By Robert C. Warner and Edith Polis

The presence of a proteolytic enzyme in milk was first reported by Babcock and Russell.² They studied the decomposition of skim milk in the presence of antiseptics and the formation of solution of the presence of antiseptics. uble nitrogenous products during the ripening of cheese and attributed the observed proteolysis to an enzyme, "galactase." Others^{3,4} have studied the same problem, and the general results of Babcock and Russell have been confirmed. No

(1) One of the laboratories of the Bureau of Agricultural and In-(1) One of the laboratories of the Bureau of Agricultural and Industrial Chemistry, Agricultural Research Administration, United States Department of Agriculture. Article not copyrighted.
(2) Babcock and Russell, Ann. Rep., Wisconsin Agr. Exp. Sta., 14, 161 (1897); Babcock, Russell and Vivian, ibid., 15, 77, 93 (1898).
(3) Van Slyke, Harding and Hart, N. Y. Agr. Exp. Sta. Bulletin 103, 215 (1901).

(4) Thatcher and Dahlberg, J. Agr. Research, 11, 437 (1917).

successful preparations of concentrates of the

enzyme have been made.

We have found that almost all of the proteolytic activity in milk is precipitated along with the casein when milk is acidified. The activity is extremely difficult to separate from casein and is absent from the usual casein preparations only if they have been exposed for long periods to alis absent from the usual casein preparations only if they have been exposed for long periods to alcohol. It is thus present in both commercial casein and most purified laboratory preparations. The fact that casein may undergo proteolysis in solution with the formation of products soluble at pH 4.6 has been noted by Robertson⁵ and Walters.⁶ However, they believed that their

(5) Robertson, J. Biol. Chem., 2, 317 (1906-1907).
(6) Walters, ibid., 11, 267 (1912); 12, 43 (1912).

ein preparations were free from enzymes and that the "autohydrolysis" was an inherent propy of casein.

The proteolysis caused by the enzyme may have considerable influence on the results of emical studies on casein, especially those deal-g with fractionation, depending on the conditions under which they are carried out. A knowledge of the presence and behavior of the enzyme is also of importance in the commercial applications of casein.

Experimental

Our attention was first drawn to this problem by the observation that a concentrated solution of commercial casein in borax rapidly decreases in viscosity with time. This effect has been noted in the literature, but no particular attention appears to have been paid to it. It is accompanied by an increase in the concentration of nitrogenous products which are soluble at pH 4.6 and is abolished by heating the solution to 80° for ten minthe changes may be the result of the activity of an enzyme which slowly hydrolyzes casein.

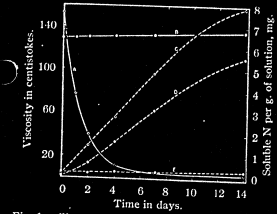


Fig. 1.—Changes in viscosity and soluble nitrogen in a casein solution at pH 8.6: A, viscosity of unheated solution; B, viscosity of heated solution; C, nitrogen soluble at pH 4.6 in unheated solution; D, nitrogen soluble in trichloroacetic acid in unheated solution; E, nitrogen soluble at pH 4.6 in heated solution.

These processes and their dependence on the pH of the solution are shown in detail by the following experiment. Reprecipitated casein prepared as previously described was dissolved in borax (0.021 g. per g. of solution) and hydrochloric acid or sodium hydroxide was added to obtain the desired pH. The casein was made up to a concentration of 9.71% by weight. The kinematic viscosity was determined in a capillary viscometer at 25° at intervals after making up the solution. The viscosity drop during the first few hours was linear with respect to time and the viscosity at zero time, i. e., at the time the alkali was added to the casein, was determined by extrapolation.

The increase in soluble nitrogen at 25° was followed by two methods. (1) A sample of one gram of the solution was weighed into a 25-ml. volumetric flask and was pre-

(7) Zoller, Ind. Eng. Chem., 12, 1171 (1920). (8) Warner, This Journal, 66, 1725 (1944).

cipitated by adding 2.5 ml. of a 1 M acetate buffer of \$\rho 14.6\$ plus sufficient hydrochloric acid to bring the sample to this \$\rho 11\$. After being made to volume, the contents were filtered, and nitrogen was determined on an aliquot of the filtrate by the method of Koch and McMeckin.\(^9\) (2) A similarly prepared sample was precipitated with trichloroacetic acid at a final concentration of 2.5%, and nitrogen was determined on the filtrate as before.

The changes in viscosity and soluble nitrogen with time are shown in Fig. 1 for an experiment at \$\rho 11\$ 8.6. The viscosity drop is rapid initially and gradually decreases in rate until it is very slow at the end of a week. The increase in soluble nitrogen continues even after the viscosity is almost constant. Data are also given for a sample of the same solution which had been heated to 80° for ten minutes immediately after the casein was dissolved. It is seen that the heated solution shows no drop in viscosity, but rather a very slow increase. This increase was followed in other cases for periods up to six months and was still continuing. The appearance of soluble nitrogen was almost completely abolished by the heating.

The influence of \$\rho\$H on the viscosity changes is shown in Fig. 2. The initial viscosity has an extremely sharp maximum at a \$\rho\$H of about 9.4. At all \$\rho\$H's below 9.8 the viscosity decreases with time. After one week it has reached its lowest value at \$\rho\$H 8.5, and this may be taken as the \$\rho\$H of optimum activity of the enzyme. At \$\rho\$H 10.3 the viscosity, although lower initially, increases with time at a rate somewhat greater than that of the heated solution at \$\rho\$H 8.6. This increase was also observed at \$\rho\$H 11.2 but was much slower. On the acid side of the optimum the viscosity drop proceeds more slowly than at \$\rho\$H 19 to 9.5, but is still evident at \$\rho\$H 5.8, the lowest \$\rho\$H and which satisfactory solutions can be prepared.

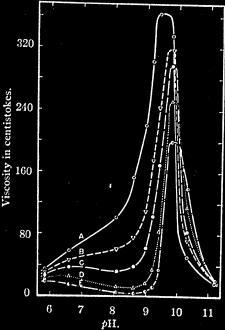


Fig. 2.—Viscosity of 9.71% casein solutions as a func tion of pH. Curves A, B, C, D and E show the viscosity at 0, 1, 2, 4, and 7 days, respectively, after the solutions were made up.

The changes in soluble nitrogen in these solutions (Fig. 3) were consistent with the viscosity data. A definite maximum was found at ρ II 8.5 with a decrease in the rate of formation of soluble nitrogen on either side of this ρ II.

⁽⁹⁾ Koch and McMeekin, ibid., 46, 2066 (1924).

The amount of nitrogen soluble in trichloroacetic acid was somewhat more than half of that soluble at \$\rho\$11 4.6 in the region of activity of the enzyme. The points on the curves at \$\rho\$14 4.6 were obtained on suspensions of casein which were allowed to stand at this \$\rho\$11 in a borate buffer in the presence of chloroform. At \$\rho\$11 11.2 the increase in the rate of formation of soluble nitrogen can be attributed to alkaline hydrolysis of the casein. After seven days there was a noticeable growth of mold at \$\rho\$11 5.8 and 6.6. The points at fourteen days in these solutions are therefore too high.

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was a noticeable growth of mold at \$11 b.8 and 0.0. The points at fourteen days in these solutions are therefore too high.

Similar experiments have been carried out on a number of samples of commercial casein. All of the results were consistent with the above interpretations, although there was a wide variation in the magnitude of both the initial viscosity and the viscosity changes with time. The variation in initial viscosity in the region of \$11 8 is probably due largely to the heat used in commercial practice in precipitating and drying the casein. By heating these dry samples further at 75° the viscosity of their solutions could be made to increase several-fold. This procedure did not abolish the decrease in viscosity with time.

Casein samples which had been exposed to ethanol at room temperature in drying contained less enzyme activity than the samples used in the above experiments in which the treatment with ethanol was of very short duration and was carried out at 2°. By treatment of a wet casein precipitate with ethanol for several hours at room temperature, particularly with concentrations around 50%, it was possible to reduce the enzyme activity to a very low level.

While there was no marked bacterial or mold growth in the solutions during the period of observation (except at \$11 5.8\$ and 6.0 as noted above) because of the bacterio static action of borax, only those solutions above \$11 9\$ were sterile. In order to eliminate bacterial activity as the cause of the casein hydrolysis, the increase in nitrogen soluble at \$11 4.0\$ was followed on a solution sterilized by filtration through a Scitz filter. The solution used had a \$14 0 sex followed on the same borax concentration but to one-half the casein concentration used in previous experiments. The filtered solution was placed in a series of sterile test-tubes and one tube was used for each determination of soluble nitrogen. The sterility of each of the samples was checked by plating with nutrient agar. Half of the solution was retained unfiltered a

COMPARISON OF SOLUBLE NITROGEN FORMED IN STERILE AND NON-STERILE SOLUTIONS

Time sampled, days	Bacteria per ml.	mfiltered sol Molds per ml.	Soluble N,a	Filtered solution ^b Soluble N, ^a
0	5300	0	1.7	1.8
2	80	20	8.6	8.1
4	10	100	18.4	18.0
7	0	130	31.4	31.5

 $^{\rm o}$ Nitrogen which is soluble at pH 4.6 expressed as percentage of the total nitrogen. b All samples of the filtered solution were sterile.

In the fractionation of casein by the imethod reported previously, some enzyme activity is found in both the α-and β-fractions after the first precipitation. In the α-fraction a small degree of activity remained even after nine precipitations, but none was present in the purified β-fraction. Only a trace of enzyme remains in whey after precipitating the casein. The enzyme was not destroyed when casein was exposed for one hour to a pH of 1.5 or 11. Chloroform was found to have an inhibitory effect on the enzyme, but none was shown by borax.

Preparation of an Enzyme Concentrate.—A number of unsuccessful attempts were made to separate the enzyme from casein. However, it was found possible to separate

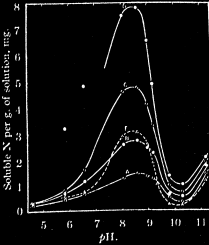


Fig. 3.— Increase in soluble nitrogen as a function of pH. Curves A, B, C and D show the nitrogen soluble at pH 4.6 at 2, 4, 7 and 14 days, respectively, after the solutions were made up. Curve E sligows the nitrogen soluble in trichloroacetic acid after seven days. Total nitrogen in all cases was 15.2 mg, per g, of solution.

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the enzyme from the hydrolysis products after allowing the enzyme to act on casein. Several concentrates were prepared by the following method:

A kilo of commercial casein was made up as a 15% solution at pl1 8.5 by dissolving in borax. Toluene was added, and the solution was allowed to stand at room temperature for two weeks. It was then dialyzed against a solution of borax for an additional two weeks. The dialyzed solution was precipitated by adjusting to pl1 4.5 with hydrochloric acid. The filtrate from this precipitation was salted out with ammonium sulfate, and the fraction appearing between 15 and 23% salt was retained. It was dissolved at a pl1 of 5.5 dialyzed, and then precipitated by adjusting the pH to 4.5. The precipitate was dissolved at pl1 5.5 to give about 25 ml. of a 0.6% solution of protein.

The various fractions of the digest were tested for enzyme activity by adding them to a solution of enzyme-free casein dissolved in borax at pl1 8.5 and following the viscosity change with time. If the sample being tested does not contain sufficient protein to increase the viscosity of the solution appreciably, the concentration of enzyme is proportional to the initial slope of the viscosity-time curve. The enzyme-free casein was prepared by heating a solution of casein to 90% for half an hour and then precipitating, washing and drying the casein. A solution of this casein showed a very slow rise in viscosity with time similar to that for the heated solution in Fig. 1.

Several of the fractions in the above separation, especially the first precipitate at pl1 4.5, showed some enzyme activity, so the fractions in the above separation in the preparation described contained more activity per gram of nitrogen than any other fraction and had on this basis about 150 times the activity of the original casein.

The enzyme appears to be insoluble at this reaction in the presence of four boundaries, the activity migr

Discussion

The proteolysis that occurs in casein solutions is attributed to the presence of an enzyme on the pasis of the following evidence: (1) The activity destroyed by heat; (2) it has a definite pH opnum; (3) the activity can be concentrated; (4) the proteolysis proceeds in sterile solutions. The enzyme is presumably the same as that studly by Babcock² and Van Slyke³ and their coworkers in their investigations on the proteolysis in skim milk and cheese. The changes observed in their experiments were very slow, and the

in their experiments were very slow, and the period of observation sometimes continued for as long as a year. However, they were carried out at a pH considerably below the optimum for the enzyme. Since the enzyme occurs in rennet casein as well as in acid-precipitated casein, it would also be present in cheese.

Babcock and Russell attempted to purify the

enzyme and found that an aqueous extract of chloroform-treated separator slime showed more proteolytic activity than milk. Separator slime contains very large numbers of bacteria and, in addition, leucocytes and other cellular debris. Chloroform will cause lysis of much of this material, and at least a part of the proteolytic activity may come from this source. Their results with this preparation have a doubtful bearing on the properties of the enzyme occurring in milk. It is not easily settled as to whether the enzyme is secreted in milk as such or whether it is of hear

is secreted in milk as such or whether it is of bacterial origin. Babcock decided in favor of the first alternative because of the invariable presence

the enzyme in milk in fairly uniform concentrations. Appreciable bacterial activity subsequent to milking was excluded in the preparation of the casein used in this work. However, bacteria are normally present in the work. teria are normally present in the udder of the cow, and, while the bacterial counts made on milk immediately after milking are usually low, these bacteria cannot be excluded a priori as the source of the enzyme.

Carpenter10 has studied the appearance of nitrogen soluble in trichloroacetic acid in solutions of casein as a function of pH. He did not give any details on the preparation of the casein used, but it is evident from his data that it did not contain any of the enzyme. Our data on soluble nitrogen by the trichloroacetic acid method at pH's 10.3 and 11.2 (Fig. 3), under which conditions the enzyme is inactive, are in fair agreement with Corportor's data in this AH range when both with Carpenter's data in this pH range, when both are calculated as percentage of the total nitrogen.

Walters reported 8% hydrolysis of casein in 137

(10) Carpenter J. Biol. Chem., 67, 647 (1926).

hours at 37.5° in a solution neutral to phenolphthalein. This hydrolysis, which is greater than that reported by Carpenter but less than that in Fig. 3, can doubtless be attributed to the enzyme.

A similar observation was made by Pertzoff. ii
The influence of pH on the initial viscosity of casein solutions, as shown in Fig. 2, is essentially in agreement with the data of Zoller. 12 He reported viscosity determinations as a function of pH in solutions of various alkalies and alkaline salts. All of his solutions exhibited maxima at a pH of about 9.2, except the ones made up with borax. The maximum at pH 8 rather than at pH 9.2, in the presence of borax, can probably be attributed to the high concentration of borax that must be used to attain a pH approaching 9. In our series of measurements at a constant concentration of borax, the viscosity maximum occurred at about the same ρ H as was found by Zoller for other alkalies. The change in initial viscosity from that at ρ H 7 to 8 to that at the maximum is considerably greater in Fig. 2, then in Zoller's considerably greater in Fig. 2 than in Zoller's

An interesting point resulting from the viscosity data was the slow increase in the viscosity of casein solutions in which the enzyme had been destroyed by heat or rendered inactive at a high pH. The effect was not studied systematically as a function of pH, but was at a maximum at pH10.3 and was much smaller in the range of pH 8 to 9 and at pH 11.3. This increase in the viscosity may be related to a slow structural change in the casein.

Acknowledgment.—The authors are indebted to Joseph Naghski for carrying out the bacterio-logical work and to T. L. McMeekin and R. W. Jackson for their advice and interest in this investigation.

Summary

Casein contains an enzyme that causes its slow hydrolysis accompanied by a decrease in the vis-cosity of its solutions and the appearance of nitrogenous products soluble at pH 4.6 and in trichloroacetic acid. The enzyme was present in all samples of commercial casein examined and in all laboratory preparations made by the usual methods.

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⁽¹¹⁾ Pertzoff, J. Gen. Physiol., 11, 239 (1927-1928).
(12) Zoller, ibid., 3, 635 (1920-1921).